UK Patent Application (19) GB (11) 2 120 666 A

- (21) Application No 8312493
- (22) Date of filing 6 May 1983
- (30) Priority data
- (31) 8215532
- (32) 27 May 1982
- (33) United Kingdom (GB)
- (43) Application published 7 Dec 1983
- (51) INT CL³ CO7C 85/12 87/28
- (52) Domestic classification C2C 22O 227 22Y 29X 29Y 311 31Y 321 326 32Y 360 361 36Y 43X 451 45Y 509 620 623 62Y 630 650 652 666 697 778 AA LJ NB NN U1S 130B 1347 C2C
- (56) Documents cited
- (58) Field of search C2C
- (71) Applicant
 Imperial Chemical
 Industries PLC
 (United Kingdom),
 Imperial Chemical House,
 Millbank, London
 SW1P 3JF
- (72) Inventor Peter John Richardson
- (74) Agent and/or Address for Service
 Malcolm John Houghton, Imperial Chemical Industries PLC, Legal Department: Patents, Thames House North, Millbank, London SW1P 4QG

(54) Hydrogenation process

(57) Hydrogenation of perhalogenated terephthalonitriles to their corresponding amines under acid conditions is suitably conducted under a pressure of 1 to 100 atmospheres and at a temperature of from 0 to 200°C in the presence of (i) a hydrogenation catalyst containing 0.1 to 70% by weight of a metal in Group 8 of the Periodic Table, (ii) an inorganic acid in an amount at least

chemically equivalent to the amine formed and (iii) a solvent which is inert to the reaction ingredients and which does not poison the catalyst; the concentration of nitrile in the total reaction mixture being from 3 to 25% by weight. Preferably water is present in the proportion of water to solvent from 1:50 to 1:1 parts by weight. The diamines obtained by this process are useful intermediates in the preparation of pesticidal compounds. 2,3,5,6-Tetrafluoroxylylene diamine and its salts are novel compounds.

GB 2 120 666 A

5

10

15

20

35

SPECIFICATION Hydrogenation process

This invention relates to a process for hydrogenating perhalogenated terephthalonitriles and to novel compounds which can be obtained therefrom.

Processes for the hydrogenation of terephthalonitrile have been numerously described and as seen from, for example, UK Patent Specifications Nos. 810530, 852972 and 1149251, the hydrogenation is normally carried out under ammoniacal conditions.

Such conditions are not altogether suitable, however, for the hydrogenation of certain halosubstituted terephthalonitriles. For instance, in the case of tetrafluoroterephthalonitrile a high molecular weight material is produced thought to be the result of nucleophilic substitution of ring fluorine atoms by an amine group of the bifunctional hydrogenation product.

According to the present invention there is provided a process for hydrogenating a perhalogenated terephthalonitrile of the formula (I):

15 in which each X is independently fluoro or chloro, to its corresponding amine of formula (II):

$$\begin{array}{c|c}
\text{CH}_2\text{NH}_2\\
\hline
\text{CH}_2\text{NH}_2
\end{array}$$
(II)

which comprises reacting the terephthalonitrile with hydrogen in the presence of a hydrogenation catalyst under acid conditions.

The diamines obtained by this process are useful intermediates in the preparation of pesticidal compounds. 2,3,5,6-Tetrafluoroxylylene diamine and salts thereof are novel compounds and form another aspect of the present invention.

Any sultable hydrogenation catalyst may be used. Generally, it will be a metal, particularly a metal in Group 8 of the Periodic Table, and will normally include rhodium, palladium, ruthenium, nickel, cobalt, platinum or copper as a component. The metal will usually be present upon a support such as carbon, alumina, alumina-silica, silica, kieselguhr, calcium carbonate, barium sulfate or bentonite. The active metal will usually be present in a proportion of from about 0.1 to 70% by weight, and, in the case of noble metals, generally 1 to 20%. A preferred catalyst is palladium preferably supported on charcoal and especially 5% palladium on charcoal. Nickel and cobalt catalysts, which tend to dissolve in acid conditions, may be less suitable.

The proportions of catalyst to nitrile may be wide-ranging. However, an amount of 0.5 to 5% by weight on nitrile has been found adequate for a catalyst containing a Group 8 noble metal.

The acid used to create the acid conditions is suitable an inorganic acid, particularly an oxyacid and ideally sulphuric acid, although other strong acids, such as hydrochloric acid, may also be suitable. At least an amount of acid chemically equivalent to the amine formed, should be used and preferably an excess up to, for instance, five times the chemically equivalent amount.

It may be prudent to add the acid continuously or intermittently during the process in case too high an acid concentration at the start of reaction, particularly when working at high nitrile concentrations, should have a deleterious effect on product yield.

5 :	It is expedient to slurry the preferably with water added. The profession the catalyst, its choiche reactant and intermediate solvents are alcohols, especially metwith respect to yield and an absert of operation, it is stirrable nitrile slurry. Suitably, and welght and even up to intermittently during the processions.	he solvent loe will be and final price will be and final price and final	should be influenced roducts an monohydra thanol. The rate at low to have suntration of eight, part	one that is in by operation dobviously in alcohols of the presence of the temperation of the color in the icularly if the local price of the color in the icularly if the	nert to the re nal consider its effect on of the formul of water give ures and pre ent and/or w total reactle a acid is add	ation, the solubility in it of yield. Particularly sultable la ROH, in which R is C ₁₋₆ es processing advantages essures. Vater present to provide a continuously or	5
15	to 1:1 parts by weight. The pressure and temper suit the hydrogenation equipm 1 to 100 or more atmospheres ranging from, for example, 0°C working at higher temperature	ature of hy ent availab c, convenie c to 200°C s to avoid (drogenation le and to a ntly from 1 and typica dehalogen	on may vary to toold too slot to 30 atmo ally from 10 ation of the p	within wide w a reaction spheres mai °C to 120°C product. For	limits and will be chosen to a. Suitable pressures of from be used at temperatures c. Care must be exercised in instance, prolonged	15
20	processing at 150°C can lead In carrying out the proce a glass lined or stainless steel solvent and water. The autock stirred at the desired temperal	to defluorings of the ingresserve is pressure until sure until sure until su	nation of to vention, the irred autoc urised to to ufficient hy	etrafluoroxyl e nitrile star clave and slu he desired e drogen has l	ylene diamii ting materia ırried with a xtent with h been absorb	ne. I is conveniently charged to If or part of the acid in the ydrogen and rotated or ed.	20
25	The amine hydrogenatio recrystallised from a water/solobtelned in its pure form by so Tetrachloroterephthalon tetrachloroterephthaloyl chlor	n product i vent mixtu Ivent extra itrile may b	s obtained re and isol ction tech e obtained	as a salt wh ated by conv niques from I from comm	lich, in the so ventional tec a strongly a nercially ava	olid phase, may be chniques. The amine may be lkaline solution of its salt. ilable	25
	which can then be dehydrated be obtained by replacement w tetrachloroterephthalonitrile. corresponding tetrachlorinate	l with, for e vith fluoro c Thus, tetrat d compoun	xample, pl of one or m luoroterep ad with pot allowing E	nosphorus of ore of the ch hthalonitrile assium fluor camples 1 to	xychloride. C hloro substit may be obt ride in a pola o 30 in whicl	Other starting materials may uents of the ained by fluorinating the ar aprotic solvent. In percentages are by weight.	30
35	Percentage yields of diamine the same molar response fact the dinitrile starting material v	ors as for t	he diamine	i. In all Exam	ipies, save 🗅	xample / (q.v.), collection of	35
40	EXAMPLE 1 Tetrafluoroterephthalon 5% palladium on carbon catal with nitrogen, and then press hours at 75°C. The resulting s solution was filtered to remov liquid surface. 74 OP Ethanol in 5N sodium hydroxide and e was recrystallised from toluer	yst (0.25 gurised with slurry was it catalyst, was added) were load hydrogen filtered and then the value of the v	ded into the to 15 atmos i the residue vater remove 06 g of a wh vith ether. T	glass liner of spheres. The slurried with ed by heat u hite solid pre he residue a	h water. The aqueous ntil a crust formed on the cipitate. This was dissolved fter evaporation of the ether	40
45	I.R. (KBr) 3385, 3275, 2955, 16	00, 1480,	1348, 126	8, 1165, 98	17, 928, 878	3, 828, 700 cm ⁻¹ .	45
	Proton nmr 2.09δ and 3.78δ , consi	stent with	(2p, s,N	IH ₂) and (2p,	, s, Ar—CH ₂	—N) respectively.	
50	UV (0.5NHCl in 50/50 methat λ max = 273 nm ϵ = 1.93 × 10 ³ λ min = 234 nm	nol/water)					50
•	Elemental Analysis	С	н	N	F		
55	Found (%)	46.5	4.0	13.6	37.1		55
_	Calculated (%) (as $C_BH_BF_4$)	46.1	3.9	13.5	36.5		- -

15

EXAMPLE 2

Tetrafluoroterephthalonitrile (5 g), methanol (70 ml), water (10 ml), 98% sulphuric acid (3.5 g) and 5% palladium on carbon catalyst (0.25 g) were loaded into the glass liner of a rotating autoclave, purged with nitrogen, then pressurised with hydrogen to 30 atmospheres. The autoclave was rotated for 5 hours, during which time the temperature rose from 15°C to 18°C, and the pressure declined to 28 atmospheres. A slurry of catalyst and solid 2,3,5,6-tetrafluoroxylylene diamine sulphate was filtered. The methanol was removed from the filtrate by reduced pressure distillation, and the aqueous residue, together with additional water, used to completely dissolve the separated solid sulphate product. 5 ml of this aqueous solution was added to 25 ml 10N sodium hydroxide solution, and extracted with four 10 ml aliquots of diethyl ether. GLC analysis of the combined aliquots showed the yield of 2,3,5,6- aminomethyl-2,3,5-tetrafluoro-benzylalcohol or 2,3,5,6-tetrafluorobenzylamine, and no 4-aminomethyl-2,3,5-tetrafluoro-benzylalcohol or 2,3,5,6-tetrafluorobenzylamine.

EXAMPLES 3 TO 6

Further tetrafluoroterephthalonitrile reductions were carried out according to Example 2, but with the autoclave charges and reaction conditions summarised in Table I. It is to be noted that the higher pressure used favoured higher diamine yields.

9/29/06, EAST Version: 2.0.3.0

Hydrogenation of tetrafluoroterephthalonitrile in rotating autoclave TABLE 1

REACTOR CHARGE	TOR CHARGE	RGE		-	S	CONDITIONS			YIEI	YIELDS	
Methanol Water ml. ml.	Water ml.		H ₂ SO ₄ 9.	Cat. g.	Temp. °C	Press. at.	Time hrs.	DA	Ą	₹	ВА
70 20	20		9.0	0.2	12—15	12—15 30—20	9	87.4	0	2.0	0.9
70 10	10	_	6.0	0.125	12	30—25	9	86.3	0.8		1.8
70 10	10		3.5	0.125	4	7—5	9	81.5	11	4.5	3.8
70 10	10		3.5	0.25	12—19	4 7	7	75.2	0.3	12.6	2.6
40 40	40		9	0.125	10	3020	9	96.0 *	0	0	0

DA = 2,3,5,6-tetrafluoroxylylene diamine CA = 4-cyano-2,3,5,6-tetrafluorobenzylamine AA = 4-aminomethyl-2,3,5,6-tetrafluorobenzyl alcohol BA = 2,3,5,6-tetrafluorobenzylamine

* conversion 61%

EXAMPLE 8

Tetrafluoroterephthalonitrile (30 g), methanol (420 ml), water (90 ml), 98% sulphuric acid (21 g) and 5% palladium on carbon catalyst (1.5 g) were loaded to a 1 litre 316 stainless steel autoclave, fitted with a glandless agitator and gas recirculation facility. The autoclave was purged with nitrogen and the contents maintained under 30 atmospheres of hydrogen pressure while agitation was continued for 6 hours. During this time the temperature rose from 20°C to 28°C. The product slurry was treated and analysed as in Example 2. The yield of 2,3,5,6-tetrafluoroxylylene diamine was 91%, with 0.3% of 2,3,5,6-tetrafluorobenzylamine.

5

EXAMPLES 9 TO 13

Further tetrafluoroterephthalonitrile reductions were carried out according to Example 8, but with the autoclave charges and reaction conditions summarised in Table IIK. Again, the higher pressures gave superior diamine yields.

0

TABLE II
Hydrogenation of tetrafluoroterephthalonitrile in stirred autoclave

Γ		Γ.,				
	BA	0.04	4	0.4	0,8	8.4
YIELDS	\$	0	1.0	1.9	1.6	3.0
YIEL	ర	0	0	0.1	0.3	3.8
	DA	88	98	83.3	75	65
60	Time hrs.	9	9	9	9	9
CONDITIONS	Press. at.	32	7	30	3.5	28
Ö	Temp.	5—15	17—15	20—18	15—18	20—15
	Cat.	1.5	0.25 17— 1.0 20— 0.25 15—	0.5		
RGE	H ₂ SO ₄	21	7	4	7	7
REACTOR CHARGE	Water ml.	06	20	9	20	70
REAC	Methanol mí.	420	280	420	280	280
	Nitrile 9.	30	10	20	10	70
	Example Nitrile No. 9.	တ	0		12	13

EXAMPLE 14

Tetrafluoroterephthalonitrile (5 g), methanol (70 ml), 98% sulphuric acid (3.5 g) and 5% palladium on carbon catalyst (0.125 g) were vigorously agitated under hydrogen at atmospheric pressure for 4.25 hours at 20°C. The resulting slurry was filtered, water added to the filtrate and the methanol removed by reduced pressure distillation. The sulphates in the residue from the hydrogenation were dissolved in the resulting aqueous solution. The products of the hydrogenation were assessed by the procedure described in Example 2. Yields were 27.4% 2,3,5,6-tetrafluoroxylylenediamine, 7.6% 4-cyano-2,3,5,6-tetrafluorobenzylamine and 9.2% 2,3,5,6-tetrafluorobenzylamine.

5

7

EXAMPLES 15 TO 25

Further tetrafluoroterephthalonitrile hydrogenations were carried out according to Example 14, but with varying catalyst and acid additions, and in some Examples, water addition. Results are described in Table III. Water is seen to enhance markedly the diamine yield.

10

TABLE III Hydrogenation of tetrafluoroterephthalonitrile at atmospheric pressure

	ВА	11.9	2.0		13.7	17.0	14.1	3.6	14.4	19.6	14.4	25.2
DS.	AA	0	2.6	0	1:1	9.0	0.7	11.7	0	4.0	10.0	6.7
YIELDS	CA	14.5	0	2.4	4.8	က	6.5	1.0	10.5	4.1	0.4	0
	ÞΦ	19.8	18	15.5	65	53	49	9	49	20	20	38.3
TIONS	Time hrs.	7.0	16.5	9	က		4.5	2.25	4	4	4	4
CONDITIONS	Temp. °C.	20	20	55	70	17	20	20	19	50	20	20
EACTOR CHARGE CONDITIONS	Cat. g.	90.0	0.5	90.0	0.25	0.125	0.25	0.25	0.125	0.125	0.125	0.125
RGE	H ₂ SO ₄	3.5	3.5	3.5	3.0	3.5	2.75	3.0	3.0	3.5	4.0	5.0
REACTOR CHARGE	Water ml.	0	0	0	S	S	5	10	0	10	5	6
REA	Methanol ml.	70	70	70	70	70	70	70	70	20	70	70
	Nitrile g.	S	Ŋ	ß	ro	က	ഹ	ъ	ĸ	ß	.c	ĸ
	Example No.	15	16	17	18	19	20	21	22	23	24	25

10

20

25

30

EXAMPLES 26, 27 and 28

Tetrafluoroterephthalonitrile (2.5 g), methanol (70 ml) and 5% palladium on carbon catalyst (0.125 g) were loaded to the glass liner of a rotating autoclave, together with 98% sulphuric acid and water in the amounts indicated in Table IV. After purging with nitrogen, the autoclave was pressurised 5 to 10 atmospheres with hydrogen, rotated and heated for the time shown in Table IV. The product suspension was filtered, and the residue and filtrate quantitatively analysed for 2,3,5,6tetrafluoroxylylene diamine by high pressure liquid chromatography.

If Examples 26 and 27 were carried out at 75°C, it could be expected that the UV spectrum of the solution phase would indicate the presence of 4-cyano-2,3,5,6-tetrafluorobenzylamine (\(\lambda\)max = 234 10 nm in 0.5 N HCI).

Examples 26 to 28 illustrate the beneficial effect of water and the better yield obtained when using a higher acid: nitrile molar ratio.

TABLE IV

Example	H,SO,:nitrile	Water	Temp.	Time	% Yield diamine			
No.	(molar ratio)	ml.	°C.	hrs.	Solution	Solid	Total	
26	1.6:1	0	90	3	48	35.3	83.3	
27	1.2:1	0	92	6	10	46	56	
28	1.2:1	2	75	6	. 8	64.1	72.1	

EXAMPLE 29

15

Tetrafluoroterephthalonitrile (10 g), 74 OP ethanol (70 ml), water (5 ml), 98% sulphuric acid (5.4 15 g) and 5% palladium on charcoal catalyst (0.5 g) were charged to the glass liner of a rotating autoclave and pressurised to 15 atmospheres with hydrogen. The autoclave was rotated at 60°C for 6 hours. The resulting slurry was filtered; high pressure liquid chromatography showed the 2,3,5,6tetrafluoroxylylene diamine yield to be 72.2% and all in the solid residue.

20 EXAMPLE 30

Tetrafluoroterephthalonitrile (2.5 g) 5% palladium on charcoal catalyst (0.125 g), sulphuric acid (6.4 g) and methanol (70 ml) were charged to a glass lined rotary autoclave. The autoclave was pressurised to 50 atmospheres with hydrogen, and rotated for 4 hours at 110°C. The resulting slurry was cooled, filtered and the solid phase recrystallised from a mixture of water and methanol. Elemental 25 analysis, fluorine nmr, proton nmr, infra red and mass spectra (the latter after the sample was heated with sodium blcarbonate), were consistent with the recrystallised material being tetrafluoroxylylene diamine sulphate.

The material gave a single peak on a high pressure liquid chromatogram, when using a mixed ion pair/electrolyte elution system.

30 EXAMPLE 31

This example is included for comparative purposes only.

Tetrafluoroterephthalonitrile (2.5 g), nickel catalyst (Harshaw 5132P) (0.8 g), methanol (70 ml) and ammonia (12 g) were charged to a glass lined rotating autoclave. After pressurising to 30 atmospheres with hydrogen, the autoclave was rotated at 110°C for 3 hours. After cooling, filtering off the catalyst, and removing residual ammonia and methanol by distillation, a brown solid remained. Infra 35 red spectroscopy indicated this to be a high molecular weight material, with some loss of ring fluorine, and the presence of an amine hydrohalide.

CLAIMS

1. A process for hydrogenating a perhalogenated terephthalonitrile of the formula (I):

40

20

25

in which each X is independently fluoro or chloro, to its corresponding amine of formula (II):

$$\begin{array}{c}
\text{CH}_2\text{NH}_2\\
\\
\text{CH}_2\text{NH}_2
\end{array}$$
(II)

which comprises reacting the terephthalonitrile with hydrogen in the presence of a hydrogenation catalyst under acid conditions.

 2. A process according to claim 1 for hydrogenating tetrafluoroterephthalonitrile to form 2,3,5,6tetrafluoroxylylene diamine.

3. A process for hydrogenating a perhalogenated terephthalonitrile of the formula (I):

in which each X is independently fluoro or chloro, to its corresponding amine of formula (II):

10

which comprises reacting the terephthalonitrile with hydrogen under a pressure of 1 to 100 atmospheres and at a temperature of from 0 to 200°C in the presence of (i) a hydrogenation catalyst containing 0.1 to 70% by weight of a metal in Group 8 of the Periodic Table, (ii) an inorganic acid in an amount at least chemically equivalent to the amine formed and (iii) a solvent which is inert to the reaction ingredients and which does not poison the catalyst; the concentration of nitrile in the total reaction mixture being from 3 to 25% by weight.

4. A process according to claim 3 in which water is present in such amount that the proportion of water to solvent is from 1:50 to 1:1 parts by weight,

5. A process according to claim 3 or 4 in which the catalyst contains from 1 to 20% by weight of a 20 metal selected from rhodium, palladium, ruthenium and platinum.

6. A process according to any one of claims 3 to 5 in which the solvent is an alcohol.
7. A process according to any one of claim 3 to 6 in which the pressure is from 1 to 30 atmospheres.

8. A process according to any one of claims 3 to 7 in which the temperature is from 10 to 120°C.

9. A process for hydrogenating tetrafluoroterephthalonitrile to form 2,3,5,6-tetrafluoroxylene diamine substantially as described with reference to any one of Examples 1 to 29.

10. 2,3,5,6-Tetrafluoroxylylene diamine and salts thereof.

Printed for Her Majesty's Stationery Office by the Courier Press, Learnington Spa. 1983. Published by the Patent Office, 25 Southampton Buildings, London, WC2A 1AY, from which copies may be obtained.